PATENT SPECIFICATION

(11)1493744

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(21) Application No. 51311/75

(22) Filed 15 Dec. 1975

(31) Convention Application Nos. 534 826 and 534 827

(32) Filed 20 Dec. 1974 in

(33) United States of America (US)

(44) Complete Specification published 30 Nov. 1977

(51) INT CL² C08L 33/00, 47/00, 67/00, 93/00

(52) Index at acceptance

C3P 102 2H4 2HY 4D3B1 4K8 8D1B 8D2A 8D2B2 8K4 8K8 D9B5AX D9C D9D2 D9D4 D9D8 D9E1 D9F1 E2 PC12X PC13A PC14B PC20B PC5 PC6A PC8B T2A T2D

C3R 3C 3D23 3N6 C14A C22 C29 C6B C6X C8R L1A L2CX L6G



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(54) SOLID PAINT

(71) We, SCM (CANADA) LIMITED, a Corporation organized and existing under the laws of the Dominion of Canada of 351 Wallace Avenue, Toronto, Province of Ontario, Canada, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

The present invention relates to a new type of paint product, namely, a solid paint having dimensional stability based on ion bonding.

Various resin compositions consisting of homopolymers and co-polymers having partially neutralized carboxylic acid groups are known. These contain between 3% and 20% of carboxylic acid residues of which less than 50 per cent of the carboxylic acid groups are neutralized with mono-valent, divalent or trivalent cations. The prior art resins, known as Ionomers, are desirable in industry because

10 the carboxylic acid groups are neutralized with mono-valent, divalent or trivalent cations. The prior art resins, known as Ionomers, are desirable in industry because they combine the utility of a thermoset polymer with the mobility and workability of the thermoplastic resin. Ionomers have lower densities than vinyl or cellulosic plastics and because of their similarity to polyethylenes find use as protective films in the food packaging industry. Ethylene-methacrylic acid co-polymers are disclosed in U.S. Patents 3,266,272, 3,338,739; and in Belgium Patents 674,595 and 600,397. Ethylene-sodium acrylate copolymers are described in Natherlands 15

PATENTS ACT 1949

SPECIFICATION NO 1493744

The following amendments were allowed under Section 29 on 5 February 1982:

Page 1, line 29, Page 2, line 14, Page 16, line 3, for millimeter read units

Page 3, line 8, delete millimeter unites insert units

Page 3, line 9, after strength. insert The Universal Penetrometer has an indicator with 0 to 380 graduations herein referred to as units.

Page 4, line 6, Page 8, line 37, Page 9, line 29, Page 13, line 39, delete mm. insert units

Page 4, line 14, delete millimeter

Page 9, Example 10 for Gel Strength (mm) read Gel Strength (units)

Page 10, Example 12 for gel strength (mm read gel strength (units)

Page 15, Example 23 for gel strength (mm.) read gel strength (units)

Page 16, line 3, after penetration insert as hereinbefore defined

Page 16, line 59, delete millimeter penetration insert units penetration as hereinbefore defined

THE PATENT OFFICE 5 March 1982

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C3P 102 2H4 2HY 4D3B1 4K8 8D1B 8D2A 8D2B2 8K4 8K8 D9B5AX D9C D9D2 D9D4 D9D8 D9E1 D9F1 E2 PC12X PC13A PC14B PC20B PC5 PC6A PC8B **T2A T2D**

C3R 3C 3D23 3N6 C14A C22 C29 C6B C6X C8R L1A L2CX L6G



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(71) We, SCM (CANADA) LIMITED, a Corporation organized and existing under the laws of the Dominion of Canada of 351 Wallace Avenue, Toronto, Province of Ontario, Canada, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

The present invention relates to a new type of paint product, namely, a solid paint having dimensional stability based on ion bonding.

Paint having dimensional stability based on ion bonding.

Various resin compositions consisting of homopolymers and co-polymers having partially neutralized carboxylic acid groups are known. These contain between 3% and 20% of carboxylic acid residues of which less than 50 per cent of the carboxylic acid groups are neutralized with mono-valent, divalent or trivalent cations. The prior art resins, known as Ionomers, are desirable in industry because they combine the utility of a thermoset polymer with the mobility and workability of the thermoplastic resin. Ionomers have lower densities than vinyl or cellulosic plastics and because of their similarity to polyethylenes find use as protective films in the food packaging industry. Ethylene-methacrylic acid co-polymers are disclosed in U.S. Patents 3,266,272, 3,338,739; and in Belgium Patents 674,595 and 600,397. Ethylene-sodium acrylate copolymers are described in Netherlands Patent 6,511,920. Many of the desirable properties of these polymers such as stress-crack resistance, transparency, grease and abrasion resistance, low permeability, high elongation, high tensile strength, and low modulus are attributed in part to a type of ionic bonding.

It has now been discovered that solid paints having effective gel properties necessary to provide dimensional stability can be prepared by cross-linking certain reactive polymers with "ion clusters" having polar molecule components. This type of ion bonding differs substantially from the solvent-free ionic bonding of the prior art compounds.

prior art compounds.

The invention provides a solid paint composition having a gel strength ranging from 100 to 200 millimeter penetration and a dimensional stability based on ion

bonding comprising the admixture of:
a. a polymer composition selected from:

(1) a solution of an ion-bonding resin having a molecular weight ranging from 1,000 to 7,000 and sufficient reactive acid functional groups selected from carboxylic, sulfonic and phosphonic groups to provide an acid number from 20 to 80, said resin dissolved in a non-polar solvent to provide a 25 to 90 weight per cent

(2) a stabilized non-aqueous dispersion of a polymer having a molecular weight ranging from 25,000 to 1,000,000 and sufficient reactive acid functional groups selected from carboxylic, sulfonic, and phosphonic groups to provide an acid number from 25 to 60, said polymer dispersed in a non-polar non-solvent as a 25 to

90 weight percent suspension; or

(3) A mixture of a non-bonding NAD resin comprising a stabilized dispersion of a polymer having a molecular weight ranging from 25,000 to 1,000,000 dispersed as a 25 to 90 weight percent suspension in a non-polar non-solvent, said resin having no



	1,493,744	2
٠	reactive functional group sites, with an ion-bonding resin solution as defined in (1), the proportion by weight of non-bonding NAD resin to ion-bonding being from 2:1 to 8:1; and	2
5	alkoxide, ammonium hydroxide, or an organic cation former dissolved or suspended in a polar solvent of high dielectric strength to provide a 10—50 weight	5
	of polymer: a mounts from 0 to 5 weight percent based on the total weight	
10	wherein said composition contains from 100 to 600 mole percent of ionic cross-linking agent per mole of acid functional group. The invention further provides a process for preparing a solid paint having dimensional stability based on ion bonding and a gel express for preparing as a solid paint having	10
15	(a) dissolving or suspending a polymer fit	
20	dimensional stability when cross-linked by ionic cross-linking agents;	15
20	suspension of an ionic cross-linking agent as defined above in a polar solvent of	20
25	The process may be followed by ageing the mixture for 3 to 25 hours at a	25
30	The invention further provides paint sticks encased in a removable skin or bars based on the above composition and processes.	
	cross-linking agent in a high dielectric polar solvent. Cross-linking of the polymers	30
35	sufficient dimensional stability under storage conditions, i.e., is self-supporting yet	35
10	For practical and protective purpose, such stick of paint will generally be	40
5	placing the paint stick in contact with the surface to be painted followed by usual vertical and lateral movements.	
	air-curable paint film is deposited thereon. The shear provided by drawing the paint stick over the surface to be painted is sufficient to cause the solid paint to deform to a flowable coating at the point of contact. Such a solid paint is one that	45
0	surface. It is assumed that the solid paint of the present invention will contain the usual pigments, fillers, driers, bonding agents, and other additives to provide films such a solid paint could be followed and hiding power. It is anticipated that	50
5	such as, for example, coil coating of metal.	55
	functional groups either built into the polymer chain or grafted thereto by the	÷
	polyesters, unsaturated polyesters, polyurethanes, polyolefins, polyacrylates, polyhydrocarbons derived from aliphatic and aromatic hydrocarbons having α,β - unsaturation, vinyl resins and chlorine-substituted vinyls as well as other	60
	combinations known to the art. Preferred are homopolymers selected from alkyd resin, polyester, unsaturated polyester, polymethacrylate and polyacrylate and	65

	copolymers thereof with vinyl chloride, an olefin or styrene or mixtures thereof, wherein the reactive acid group is a carboxylic acid group. The particular reactants and quantities are chosen to produce resins having pendant and/or terminal functional substituents which are capable of further reaction with ionic	
5	reagents to form gels of proper dimensional stability and gel strength. Desirable application properties result when the gel strength is from 100 to 190 and preferably from 135 to 180 when measured 25 hours after gelling. Gel strength is recorded in millimeter unites using a Universal Penetrometer — the lower the penotrometer reading, the higher the gel strength.	5
10	It is essential that the resin as shown in a(1) above be soluble in a non-polar solvent and that the resin having pendant and/or terminal functional reactive groups which are readily ionizable. Such ionizable groups are the anionic functional sulfonic, phosphonic and carboxylic groups. The carboxylic acid functionality is especially preferred since a variety of polymers having such	10
15	reactive ionizable groups can be readily purchased or synthesized. Preferred reaction products are those obtained from the combination of carboxylic acid substituted polyesters and alkyd polyesters having molecular weights in the range from 1000 to 7000 which contain from I to 4 reactive functional groups per each 2.000 unit of molecular weight. Polyesters and polyethers having molecular	15
20	weights in the range of 400—2000 and which yield solid paints of desirable gel properties are especially preferred. Most preferred resins are polyester alkyd resins having a molecular weight of 1500 to 3500 and an acid number of 38 to 48 which is reacted with 210—250 mole percent of sodium hydroxide as a 25 weight percent solution basis mole of carboxylic acid group. Alkyd resins modified with	20
25	fatty acid groups and having terminal carboxylic functionality are exemplified in the best mode Examples. In the case of polyolefins, polyacrylates and other systems where no air-curing will occur, a higher molecular weight of the order of 100,000 is usually necessary. However, 1 to 4 reactive functional groups are still required per 2,000 unit of molecular weight. The alkyd resins useful in the practice	25
30	of this invention are prepared by polymerizing the polymer monomers and other intermediates in a fusion cook at a temperature of $400-600^{\circ}$ F. to yield resins having an acid value (A.V.) ranging from 30 to 55 and preferably 41 ± 2 . Certain 'longer' oil resins as exemplified hereafter in Examples 1 and 2 are polymerized at 450° F. to an A.V. of 43.0 . A preferred composition according to the invention is	30
35	one wherein a polyester resin having a molecular weight of 1500—3500 and an acid number of 38—48 is dissolved in mineral solids and wherein sodium hydroxide is present at 220—280 mole % excess basis the acid functional group. The above described polymers having ionizable reactive groups are dissolved in sufficient non-polar solvents to provide solutions having non-volatile (N.V.)	35
40	contents of from 10 to 90 and preferably from 35 to 60 weight percent. Especially preferred are solutions of 50% N.V. Suitable non-polar solvents for dissolving the polymer include both aromatic and aliphatic-type hydrocarbons and are selected based on the particular resin used, the functionality on said resin, and the nature of ionic reactant. In general, suitable solvents are hydrocarbons having a boiling	40
45	point of 125 to 400° F. and which contain up to twelve carbon atoms. These include hexane, heptane, octane, nonane, decane and mixtures thereof. Preferred hydrocarbons are the various octanes because of their suitable evaporation rates. Mineral spirits is an especially preferred solvent because of its availability and the desirable properties of the resultant solid paint. In certain cases aromatic	45
50	hydrocarbons such as toluene and xylene can advantageously be used and are especially valuable in dissolving the higher molecular polymers. It is understood that the solvent, resin and proportions of each will vary and depend on the type of resins, the type of solvent, the fillers and the other additives needed for a particular end-product solid paint. The additives, driers and other	. 50
55	usual dispersant aids can be blended with the resin solution using a Cowles agitator. The order of addition is usually not critical. If desired, the pigments and other additives may be blended with the resin material prior to the solution of the resin in the non-polar solvent. After the additives are thoroughly mixed, the resulting composition is advantageously allowed to age for 12 to 20 hours before	55
60	reacting with the ionic component. In the stabilized dispersion of a polymer in a non-polar non-solvent medium as shown in a(2) above, the polymers useful in the present invention include homopolymers and copolymers and mixtures thereof having appropriate	60
65	functional groups either built into the polymer chain or grafted thereto by the usual graft techniques. Such resins include but are not limited to polyethers,	65

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5	unsaturated polyesters, polyurethanes, polyacrylates, vinyl resin and chlorine- substituted vinyls as well as other combinations known to the art. The particular reactants and quantities are chosen to produce a resin having pendant functional substituents which are capable of further reactions with ionic reagents to form gels of proper dimensional stability and gel strength. Desirable application properties result when the gel strength is from 130 to 210 and perfolation properties	
	when measured 25 hours after gelling. A preferred composition comprises a non-aqueous dispersion of a resin having a molecular weight of 100,000 to 300,00, an acid number of 30—40, and suspended in minoral spirits.	5
10	resin is a vinyl acetate-ethyl acrylate methacrylic acid terpolymer having an acid number of 25 to 80 and is reacted with 210—250 mole per cent of sodium hydroxide as a 25 weight per cent solution in action of sodium	10
15	Gel strength is recorded in millimeter units using a Universal penetrometer— the lower the penetrometer reading, the higher the gel strength. For the a(2) type of resin formulation used in the practice of this invention, it is essential that the particular resin be insoluble or only lightly swelled by the non- solvent, as is necessary for any non-aqueous dispersion, and that the resin have pendant reactive groups which are readily ionizable. The ionizable groups are the anionic sulfonic, phosphonic and carbovelies.	15
20 25	functionality is especially preferred since a variety of polymers having such reactive ionizable groups can be readily purchased or synthesized. Preferred resins are copolymers of unsaturated hydrocarbons and unsaturated acids having molecular weights in the range of 100 000 to 200 and unsaturated acids having	20
	particle size range of from 0.01 to 30 microns. Acrylate and methacrylate copolymers having terminal carboxy functionality are especially preferred and are illustrated in the best mode examples. Non-aqueous dispersions (NADs) known to	25
30	1, 1973 at pages 14—28. The polymer exemplified in a(2) is required to have ionizable reactive sites whereas the non-bonding NAD resin of a(3) does not have such active sites. The a(3) mixtures contain a non-bonding (no reactive sites) NAD resin in combination with an a(1) resin which we have	30
35	resin. In the formation of such NADs the choice and level of stabilizer is eminently important to provide solid paints having desirable application flow and coalescence characteristics. Other useful NAD resin include those derived from polyacrylate and polymethacrylate resin and emission include those derived from	35
40	acetate, hydroxyethyl acrylate and 2-hydroxypropyl methacrylate. The polymer resins useful in this aspect of the invention can be prepared by solution polymerization followed by dispersion in a non-solvent or by dispersion polymerization. The first method invention in a non-solvent or by dispersion	40
45	ranging from 20 to 80 and preferably from 25 to 60. The second and preferred intermediates in a non-solvent under free radical conditions at a ranging from 20 to 80 and preferably from 25 to 60. The second and preferred intermediates in a non-solvent under free radical conditions at a	45
50	The above described polymers having ionizable reactive groups are dispersed in a non-polar non-solvent to provide a dispersion having non-volatile (N/V) preferably a stabilizer is present the polymers.	50
55	acid value in the range of 25 to 60.1. Especially preferred are dispersions of hydrocarbons which are selected based on the particular resin, the functionality on said resin and the nature of the initial control of the functionality on	55
60	said resin and the nature of the ionic reactant. In general suitable non-solvents are hydrocarbons having a boiling point of 100 to 400°F, and which contain up to twelve carbon atoms. These include hexane, heptane, octane, nonane, dodecane and mixtures thereof. Preferred hydrocarbons are the various octanes because of their suitable evaporation rates. Mineral spirits is an especially preferred solvent because of its availability and the desirable properties of the resultant solid paint.	60

For some resin systems aromatic hydrocarbons such as toluene or xylene may be used. It is recognized that NAD resins can be suitably formulated with various stabilizers known to the art. The function of these stabilizers is primarily to prevent the resin particle from coalescing on storage and during formulation into solid paint products. Useful stabilizers include those described and referenced in the above noted article by Dowbenko and Hart, Polyene stabilizers, which are useful for certain solid paint compositions include low molecular weight polybutadiene, grafted to a backbone of an acrylic copolymer. For the instant solid paints NAD resin stabilized by copolymers of methyl methacrylate and further reacted with 12-hydroxystearic acid and/or poly (lauryl methacrylate) are especially preferred. It is additives needed for a particular end-product solid paint. The additives, dries and other usual dispersant aids are preferably blended with the resin dispersion using a Cowles agitator. The order of addition is usually not critical. The typical solid paint formulations as described herein are of the latex type non-aqueous resin dispersion and do not usually require specific drier components to give suitable film properties; when driers are added they are used in quantities less than 2 percent and preferably less than 1 percent per weight of total composition. The driers are added for the small amount of oil or alkyd that is normally added to the film. After application the resin particles coalesce and fuse to give a did hydrocarbon), the formulation of a(2) (stabilized dispersion in a non-polar hydrocarbon), the formulation of a(3) (mixture of one). The polymer formulations shown in a(1) (solution of polymer in a non-polar hydrocarbon), the formulation of a(2) (stabilized dispersion to lon-bonding resin a(1) with a non-bonding NAD resin) are next combined with the inonic cross-linking agents of continual to a subject of the properties of the polar solvent molecules, act as reversible cross-links to join the r			
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dispersion and do not usually require specific drier components to give suitable film properties; when driers are added they are used in quantities less than 2 percent and preferably less than 1 percent per weight of total composition. The driers are added for the small amount of oil or alkyd that is normally added to the formulation to aid dispersion of the pigment and to aid in the coalescence of the film. After application the resin particles coalesce and fuse to give a dry film in matter of minutes. The polymer formulations shown in a(1) (solution of polymer in a non-polar hydrocarbon), the formulation of a(2) (stabilized dispersion in a non-polar non-solvent) or the mixture of a(3) (mixture of ion-bonding resin a(1) with a non-bonding NAD resin) are next combined with the ionic cross-linking agents of dissolved in a high dielectric polar solvent. The mixture of a(3) preferably is in a proportion by weight of non-bonding non-aqueous dispersion to ion-bonding resin solution of from 2:1 to 4:1. Suitable ionic cross-linking reactants include inorganic salts which produce on solution specific cations or anions capable of combining with the terminal reactive groups of the resin to form ion clusters responsible for gel formation. Such clusters, which contain the high dielectric polar solvent molecules, act as reversible cross-links to join the reactive resin molecules in webs thus imparting gel strength and dimensional stability to the resultant solid paint. When the reactive terminal sites on the polymer are carboxylic acid groups (—COOH), the preferred cross-linking reactants include the oxides and hydroxides of sodium, potassium, lithium, barium, calcium, manganese and magnesium, the hydroxides being the preferred reactants. Equally effective cross-linking agents are the corresponding metal alkoxides it. e. sodium methylate. Additionally, ammonium hydroxide and organic cation formers such as tetramethyl-ammonium hydroxide can be used as cross-linking gelation derived by reacting sodium hydroxide can be used	15	additives needed for a particular end-product solid paint. The additives, driers and other usual dispersant aids are preferably blended with the resin dispersion using a Cowles agitator. The order of addition is usually not critical. The typical solid	15
125 matter of minutes. The polymer formulations shown in a(1) (solution of polymer in a non-polar hydrocarbon), the formulation of a(2) (stabilized dispersion in a non-polar non-solvent) or the mixture of a(3) (mixture of ion-bonding resin a(1) with a non-bonding NAD resin) are next combined with the ionic cross-linking agents dissolved in a high dielectric polar solvent. The mixture of a(3) referably is in a proportion by weight of non-bonding non-aqueous dispersion to ion-bonding resin solution of from 2:1 to 4:1. Suitable ionic cross-linking reactants include inorganic salts which produce on solution specific cations or anions capable of combining with the terminal reactive groups of the resin to form ion clusters responsible for gel formation. Such clusters, which contain the high dielectric polar solvent molecules, act as reversible cross-linking to reactive resin molecules in webs thus imparting gel strength and dimensional stability to the resultant solid paint. When the reactive terminal sites on the polymer are carboxylic acid groups (—COOH), the preferred cross-linking reactants include the oxides and hydroxides of sodium, potassium, lithium, barium, calcium, manganese and magnesium, the hydroxides being the preferred reactants. Equally effective cross-linking agents are the corresponding metal alkoxides i.e. sodium methylare. Additionally, ammonium hydroxide and organic cation formers such as tetramethyl-ammonium hydroxide and below. The cross-linking gelation derived by reacting sodium hydroxide with the above described resin molecules having terminal or pendant carboxyl groups is especially preferred. Suitable gels result when an effective amount of the cationic base combines with the free carboxylic acid functionality. In every case an amount of base substantially in excess of the amount required for neutralization is necessary to be effective. By substantial excess is meant from 100—600 mole percent of ionic reactant dissolved in the polar solvent. Although the amount of excess varies with each	20	dispersion and do not usually require specific drier components to give suitable film properties; when driers are added they are used in quantities less than 2 percent and preferably less than 1 percent per weight of total composition. The driers are added for the small amount of oil or alkyd that is normally added to the	20
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	65	the molar content of the reactive functional group i.e. moles free COOH.	. 65

	generally those solvents having a dielectric constant greater than 10, include aliphatic alcohols containing one to ten carbon atoms and one to two hydroxy groups. Although C ₁₋₈ aliphatic alcohols are usually preferred, glycols containing the similar carbon chains are sometimes useful in producing desirable gel	
5	properties in the resultant solid paint. Useful alcohols include methanol, ethanol, isopropanol, n-propanol, the normal and isomeric butanols, pentanols, hexanols, heptanols, octanols, as well as the corresponding glycols derived therefrom. Methanol is the preferred alcohol because of its costs, availability and the favorable solubility of the ionic reagents therein and is preferably used with	5
10	sodium hydroxide as the cross-linking agent. In certain applications it is preferred to use glycols or mixtures of glycols and alcohols as the plasticizer carrier for the ionic reactant. Preferred glycols are ethylene glycol and propylene glycol although for certain resins the higher glycols such as pentanediol and hexanediol act in the nature of a plasticizer and provide desirable jubricity. Additional high dielectric	10
	polar solvents useful in the practice of this invention include, water, formamide, dimethylformamide, and dimethylsulfoxide. The metal driers suitable for the instant solid paint compositions are those known to the art and include the metal salts and/or esters of various organic carboxylic acids containing up to 30 carbon atoms and mixtures thereof. The metal	15
20	salts of cobalt, zinc, zirconium, magnesium, aluminum and manganese prepared from branched chain C_{8-12} carboxylic acids are preferred driers, preferably used in an amount from 1 to 3 weight percent. Some paint formulations especially those exemplified by a(1) above require the addition of a metal drier in an amount in the order of 0.5 to 5 percent based on the weight of the resin. Whether a drier is needed	20
25	and the amount thereof depends to some extent on the oil or other source of double bonds used in the paint system i.e. number and type of double bonds available. Examples of cation formers are (1) primary, secondary, tertiary and cyclic amines, which react with hydrogen halides and hydrogen halides to give	25
30	quaternary halides to give quaternary salts; (2) substituted phosphines which combine with halides to give phosphonium salts; (2) sulfides which react with alkyl halides give sulfonium salts; (4) cyclic ethers which react with acids give oxonium salts. For industrial coating purposes, the block of solid paint is advantageously	30
35	contained in conventional holding and applicator devices. Such devices, which will vary with the nature of the substrate to be coated and will be adaptable to continuous application, usually include a device for holding the solid paint and a mechanism for adjusting the pressure placed on the paint block to allow proper deformation to provide a fluid coating and film of required thickness. Increasing	35
40	the pressure applied to the solid paint will result in the deposit of a heavier coating. Although the instant solid paints are capable of air drying, it is contemplated that for industrial coating applications curing of the film may be accelerated by the use of heat, and other energy techniques known to the art. The following specific examples illustrate only a limited number of embodi-	40
45	ments; accordingly, the invention is not limited thereto. All parts and percentages being by weight unless otherwise indicated. The driers used were commercially available conventional driers. The "mineral spirits" and the "odorless mineral spirits" had a boiling range of 300—400°F. and 345—410°F. respectively. Molecular weights reported are number average molecular weights unless otherwise specified.	45
50	Examples 1 to 12 exemplify the a(1) type of polymer solution formulations whereas Examples 13 to 19 exemplify the a(2) type polymer dispersion formulations. Examples 20 to 23 exemplify mixtures of the a(1) type of polymer solution formulations used in conjunction with non-bonding (no reactive acid functionality) non-aqueous dispersions.	50
55	EXAMPLE 1.	55
	Resin A was prepared by polymerizing a mixture (in amounts shown below) of trimethylolethane (TME), dehydrated castor fatty acid (DCOFA), Azelaic dimer acid (AZELAIC 1110) and dimer acid (EMPOL (Trade mark) 1014! at 460°F, as a fusion cook to an acid value of 41 (41 + 2 normal range).	€
60	Resin B, a "longer" oil resin, was prepared in a fashion similar to Resin A by polymerizing at 450°F, to an acid value of 42.0. Resin C, prepared using Pentaerythritol (PE) in place of trimethylolethane (TME), was polymerized at 460°F, to an acid value of 42.0. Resin D, prepared using a combination of DCOFA and Tung Oil instead of	60
65	simply DCOFA, was polymerized at 460°F, to an acid value of 43.0.	65

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	Material	TABLE I. Mols	Wt.	Acid Value
Resin A	TME DCOFA AZELAIC 1110 EMPOL 1014	2.46 2.46 1.78 0.74	295 690 340 423	41
Resin B	TM5 DCOFA AZELAIC 1110 EMPOL 1014	2.0 2.4 1.42 0.59	240 672 270 337	43
Resin C	PE DCOFA AZELAIC 1110 EMPOL 1014	1.0 2.0 0.72 0.29	136 560 135 168	42
Resin D	TME DCOFA TUNG OIL AZELAIC 1110 EMPOL 1014	1.0 0.6 0.19 0.48 0.97	120 168 168.5 91.6 555	43

EXAMPLE 2.

The polyester Resin A (25 parts) was formulated into a hydrocarbon solution by mixing with 12 parts tung oil, 13 parts mineral spirits, 2.0 parts of a cobalt drier (12.0 percent metal), 2.0 parts manganese drier (9.0 percent metal) and 3.5 parts zirconium drier (12.0 percent metal) and the resultant composition was allowed to mature at room temperature for 16 hours. Titanium dioxide (40 parts) and calcium carbonate (10 parts) were blended with the resin solution under Cowles agitation to yield a #6 Hegman grind. Various weights of sodium hydroxide were then added as a 25 weight percent solution in methyl alcohol to form the solid paints identified in Table II. Solid Paint 2A exhibited a streaky film appearance, the paint was slightly too hard requiring too much effort to apply i.e. exhibited too much drag on application, and application characteristics which were too hard. The solid paints 2B and 2C with gel strength of 147 and 161 respectively exhibited satisfactory application characteristics and film appearance i.e. the paint didn't require too much force to apply and the resultant film was uniform. All three solid paints exhibited dimensional stability and gave a satisfactory dry coating on application to a test panel surface. to a test panel surface.

Exp. No.	Resin	TABLE II. Parts NaOH Added	Percent Neutralization Calculated on Moles Carboxylic Acid	Gel Strength* (24 hours)
2A	A	6.65	225	119
2B	A	6.35	215	147
2C	A	6.05	205	161
3A	C	7.2	240	160
3B	C	7.2	240	155
3C	C	6.0	200	178
4A	C	4.75	160	176
4B	C	5.0	170	138
8A	C	6.1	200	165
8B	C	6.5	220	155

^{*}Average of three determinations

	1,493,744	8
5	EXAMPLE 3. Resin C was formulated into paints 3A and 3B using the procedure outlined in Example 2 and the same relative amounts of resin, tung oil, mineral spirits, cobalt drier, manganese drier, zirconium drier, titanium dioxide, and calcium carbonate. A third paint formulation 3C was similarly prepared from Resin C but contained 1.3 parts of cobalt drier (12% metal), 0.5 parts manganese drier (9.0% metal), 3.0 parts zirconium drier (12% metal) and 0.19 parts aluminum stearate. The solid paints formed on the addition of 25% methanolic sodium hydroxide identified as 3A, 3B and 3C each exhibited satisfactory gel strengths, application characteristics, film appearance and drying quality.	5
	EXAMPLE 4.	
15	mixing with 12 parts tung oil, 13 parts mineral spirits, 0.95 parts cobalt drier and 2.1 parts zinc drier (16 percent metal). A second resin formulation for Resin C was identical to the above except it contained only 0.9 parts of cobalt drier and additionally contained 0.45 parts of manganese drier. These resins and paints made therefrom which contained 50 parts titanium dioxide and no calcium carbonate are identified respectively as 44 and 48 in Table 18.	15
20	with neutralization values of 160 and 170 exhibit gel strengths of 176 and 138 respectively. The application characteristics of 4A were slightly inferior, the solid paint tended to be too soft. The film appearance and drying quality of both paints were acceptable.	20
25	EXAMPLE 5. Repeating the experiments 2A, 2B, 3A, 3B and 3C but adding the driers subsequent to the addition of pigment to the resin will result in essentially similar acceptable gel strengths, application characteristics and drying rates.	25
30	Paint blocks of approximate size 4" × 6" were stored using a thin SARAN (Trademark of the Dow Chemical Company) envelope for a period of six months. Application of these paints to a test panel after the storage period showed no detectable deterioration of the application and film characteristics. Additionally, solid paints prepared from the same resins but having acid values in the range of obtained when oiticica fatty acid, safflower fatty acid, soya fatty acid, or linseed fatty acid was used instead of dehydrated castor oil fatty acid.	30
35	fatty acid was used instead of dehydrated castor oil fatty acid. The best application properties were obtained when the gel strength as measured by the Universal penetrometer was between 130 and 180 mm. Gel strengths of from 100 to 130 and 180—190 gave effective solid paints with somewhat less desirable characteristics.	35
40	Resin D was prepared by first esterifying the dehydrated castor fatty acid (168 parts) with trimethylolethane (120 parts) at a temperature ranging up to 480°F, to further reaction with a tung oil (168.5) in the presence of 2.0 parts of litharge catalyst until the product was completely miscilla in the presence of 2.0 parts of litharge	40
45	product was combined with Azelaic 1110 (91.6 parts) and Empol 1014 (555 parts) and cooked to an acid value of 43.0. The resulting resin had an approximate	45
50	A cationic Resin E was prepared by condensing Resin D (1040.4 parts) with N,N-diethylaminoethanol in the presence of litharge (2.0 parts) catalyst using reaction conditions such that the predominant reaction was esterification rather than amide formation. After removal of water and excess N,N-diethylaminoethanol, Resin E had a molecular weight of 1500.	50
55	Gelation of Resin E was effected by neutralizing (100 and 300°) a 50/50 weight percent solution of Resin E in mineral spirits with 37% hydrochloric acid. The resultant solid paints had properties inferior to those of a corresponding gel neutralized to 200 percent with 32 N-sulfuric acid and resulting in gel strengths of from 100—150.	55
60	Polyester resin C (25 parts) was formulated into a hydrocarbon solution by mixing with 12 parts tung oil, 13 parts mineral spirits, 0.6 parts cobalt drier (12.0 percent metal), 0.6 parts manganese drier (9.0 percent metal) and 6.0 parts	60

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zirconium drier (12.0 percent metal) and the resultant composition was allowed to mature at room temperature for 16 hours. Titanium dioxide (40 parts) and calcium mature at room temperature for 10 hours. Intantum Gloxide (40 parts) and calcium carbonate (10 parts) were blended with the resin solution under Cowles agitation to yield a #6 Hegman grind. Various weights of sodium hydroxide were then added as a 25 weight percent solution in methanol under reduced pressure in a 'vacuum Cowles' to form a solid paint (Table II). This manner of addition diminishes the chance of entrapping air into the 'final' solid paint. Paints 8A and 8B (of Table II) exhibited superior film appearance and application properties. Both paints were dimensionally stable and exhibited good dry on application to a test panel surface. dimensionally stable and exhibited good dry on application to a test panel surface.

EXAMPLE 9.

Resin F was prepared under free radical conditions as follows: 10 parts methacrylic acid, 90 parts lauryl methacrylate, 1 part Bis (4-t-butylcyclohexyl) metnacrync acid, 90 parts lauryl metnacrylate, 1 part bis (4-t-butyleyclonexyl) peroxycarbonate (initiator), and 300 parts mineral spirits were added to the kettle. Polymerization was accomplished by heating to 60°C. and holding at this temperature for 2 hours while the mass in the kettle was being agitated. Conversion of 99% was achieved; acid value of the polymer was 65.0. Approximately 100 parts of the mineral spirits were removed by vacuum distillation.

Various weights of sodium hydroxide were added as a 25 weight percent solution in methanol to 75 parts of the 33 percent N/V resin with agitation as

shown:

Exp. No.	Parts NaOH Added	Percent Neutralization Calculated Of Carboxylic Acid		
A	6.9	150		
В	9.2	200		

The two 'clear' paints can be described as follows: Experiment A resulted in a product that was just barely dimensionally stable and exhibited poor application characteristics, i.e. on applying the paint laid down too thick a film and too much force (relative to the previous examples) was required to draw the sample across the test panel.

Experiment B resulted in a stronger product that exhibited good dimensional stability (gel strength of approximately 160 mm penetration) and good application characteristics. Paint B exhibited very little drag on application. Both these products resulted in a 'dry' film on the test panel.

EXAMPLE 10.

Resin G, a 100 percent N/V dicarboxypolybutadiene having a molecular weight of 1410 and an acid value of 65.0, was formulated into the following solid paint systems:

paine systems.				
Exp. No.	A	В	С	D
Resin G (parts)	50	50	50	17
Resin A				33
Mineral spirits	50	50	50	50
Cobalt drier (12 percent metal)	.5	.5	.5	.5
Zirconium drier (12 percent metal)	1.7	1.7	1.7	1.7
Titanium dioxide	-	130	110	90
Calcium carbonate	-	70	5 0	40
NaOH (25 parts in methanol)	18.0	20.2	36	24
Percent Neutralization	200	300	400	350
Gel Strength (mm.)	250	180	110	160

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Paint A having a gel strength of 250 did not exhibit dimensional stability. Paints B, C and D were dimensionally stable. Under application action Paint B tended to put down too thick a film and was a little too elastic, i.e. tended to be slightly taffy like. Paint C was too hard and for this reason it resulted in poor quality application. Paint D exhibited dimensional stability and acceptable application. All the paints resulted in a dry film on the test panel.

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Alkyd Resin H was prepared by polymerizing a mixture of 146 parts trimethylolpropane, 146 parts pentaerythritol, 908 parts dehydrated castor oil fatty acid, and 413 parts Azelaic dimer acid (AZELAIC 1110) at 480°F, as a fusion cook to an acid value of 42. The resulting resin exhibited a viscosity of Z₂ as determined using the Gardner-Holt Bubble Tube Test method ASTM D1545.

Alkyd Resin I was prepared by polymerizing a mixture of 116.5 parts

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Alkyd Resin I was prepared by polymerizing a mixture of 116.5 parts trimethylolpropane, 116.5 parts pentaerythritol, 296 parts dehydrated castor oil fatty acid, and 821 parts Azelaic dimer acid (AZELAIC 1110) at a temperature of 460°F to an acid value of 30. The resulting resin exhibited a viscosity of Z₂+

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EXAMPLE 12. Solid paints were prepared from Resins H and I according to the procedure of Example 2 with the exception that driers were allowed to mature at room temperature for 1 hour, the order of addition of ingredients being as given in the following table with blending to a #51 Hegman grind.

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Material	Parts				
Experiment No.	1	2	3	4	
Resin I	-			50	
Resin H	50	50	37	30.	
AC 100°)	30	30	54	30	
Dramatone Blue tinting baseb	2.5	_	_		
Titanium Dioxide	100	100	100	100	
Min-u-Sil 10	10	10	10	10	
Celite 499 ^a	10	10	10	. 10	
Rheox 1°	1.0		1.0	1.0	
Odourless mineral spirits	50	55	44	50	
Cobalt drier (120 percent metal)	0.3	0.3	0.3	0.3	
Manganese drier (9.0 percent metal)	0.15	0.15	0.15	0.15	
Zirconium drier (12.0 percent metal)	3.0	3.0	3.0	3.0	
Methyl ethyl Ketoxime	0.2		0.2	_	
sodium hydroxide — methanol (24 percent sodium hydroxide)	8.0	8.0	8.0	8.0	
oneutralization .	130	130	180	160	
gel strength (mm	170	170	160	180	

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Notes to table.

a) a diluent alkyd resin not capable of direct participation in ionic bonding—Reichhold Chemicals (Canada) Ltd. DRAMATONE is trademarked product of GLIDDEN-DURKEE, Division of SCM Corporation

c) Crystalline silica product of Pennsylvania Class Sand Corp.
d) Trade mark for diatomaceous silica product of Johns-Manville Co.

e) Bodying agent product of N. L. Industries. The solid paints 1, 2, and 3 exhibited dimensional stability and characteristics equivalent or superior to the solid paint products of the previous Examples. When applied to a substrate by contact and hand pressure desirable surface films were obtained which are air cured overnight.

PREPARATION OF NAD RESINS.

The NAD resins 1, 2, 2A, 3 and 4 were prepared by addition polymerization of various monomers in the presence of non-solvents, free radical initiators and various stabilizers in the relative proportions shown in Table III. A small portion of various stabilizers in the relative proportions shown in Table III. A small portion of the monomers is charged to the polymerization kettle with the non-solvent and about 50 percent of the desired stabilizer and polymerization is initiated by heating to a reflux temperature in the order of 70—80°C. Thereafter the remaining monomers, stabilizer (30%) and free radical initiator are added with ethyl acrylate in one feed stream while the acidic component i.e. methacrylic acid and remaining stabilizer (20%) is added in a separate feed stream over a two to three hour addition period at the reflux temperature. Additional initiator (‡ total amount) is introduced in ethyl acetate in two portions over a further reaction period of 2 hours. After refluxing for an additional two hours, low boiling solvent is removed by heating to approximately 90°C. For this present invention it is important that the NAD be prepared with the carboxylic sites (or other ionizable sites) at the surface of the particle (or at least the majority be available to the surface) in order surface of the particle (or at least the majority be available to the surface) in order to provide the external acid sites on the suspended polymer particles. In this case the acid feed was started 10 minutes after the other monomer feed was commenced; and the acid feed was completed approximately 10 minutes after the other monomer feed was terminated.

Variants of the conditions shown in this example may be used as long as a

stable NAD is produced where the acid sites are available for gelling and not buried in the body of the particle. It is recommended that an acid value determination be made on the NAD.

TABLE III — GENERAL MAKEUP OF NAD POLYMERS (PARTS BY WEIGHTS)

	NAD-1	NAD-2	NAD-2A	NAD-3	NAD-4
Vinyl Acetate	227	142	142	142	142
Ethyl Acrylate	104	212	237	237	212
NAD Stabilizer	76.4	7 6	76	57.5	58
Methacrylic Acid	28	26	38	30	18
Mineral Spirits	300	300	300	300	300
Hexane	300	300	300	300	300
Azobisisobutyronitrile	7.5	7.:	5 7.5	7.5	. 6
Ethylacetate	25	25	25	25	25
Non-Volatile Content (final)	42.4	47.	5 49	49.8	58.1
Acid Value NAD	44.7	39.	5 55.5	43.7	28.5

			1,4	93,744	•	12
5	reaction wa had an acid under nitro ketone and value of 4.3 off at the polymerized the present	e were heat is monitored value of 34, gen with 82. 10 parts tri and a non-vend of the lunder free	ed together at 1 by collecting the 2 (calculated 33 3 parts glycidyl ethylamine to y olatile content of reaction). The radical condition	F NAD STABILIZER. 3.5 parts tetraisopropy 200°C under a nitroger te by-product water. Th b. This product was furth methacrylate using 400 teld a second intermed f 93.4. (The methyl ethy is second intermediate ons with 225 parts meth trts), dodecyl mercapt dical initiator. The stab	I titananate and 60 atmosphere. The resulting product her reacted at 90°C parts methyl ethyl iate having an acid l ketone is stripped (321 parts) was ayl methacrylate in	5
15	1110 dimer a	rythritol, 56 cid, and 168 alkyd resin 1	0 parts dehydra	ALKYD MODIFIER olymer was prepared be ted castor oil fatty acid 14 dimer acid in a fusion arboxylic acid function	by condensing 136 , 13f parts Azelaic	15
20	120 parts tita	mium diori	parts of 50 N to a #6 Hegman	PLE 13. I/V suspension in mir grind with 30 parts of a green used in the formula view of the second	ikyd modifier and	20
25	parts alkyd n hydroxide (23 identified in respective se	nodifier and 5% solution Table IV as	i 115 parts titan in methanol) w s Experiment IA	ium dioxide. Various vere then added to form A, and IB. Solid paints	s blended with 25 veights of sodium n the solid paints if IA and IB with	25
	test panel. By paint across the work requirements for a thir	y good appl he test pand uired to acc	ication characted a uniform film complish this is	ristics it is implied that of paint is transferred not excessive.	n application to a t on drawing the to the panel and	30
5	zirconium dri 16.1 parts sod the solid pain dimensional	ent cobalt), er (12 perce ium hydroxi t identified stability ha	0.5 parts mangent metal); the cide (25% solution in Table IV as	anese drier (8 percent driers are added for the in methanol) were the experiment IC. This p	metal), 4.0 parts alkyd modifier. added to form	35
0	excellent dry	on applicati	on to a test par	nel.	nd exhibited an	40
	Exp. No.	Resin	TABLI Parts NaOH	Percent Neutralization	Gel Strength*	•

-	Exp. No.	Resin	Parts NaOH Added	Percent Neutralization Calculated on Moles Carboxylic Acid	Gel Strength* (24 hours)
	IA	NAD-2	15.8	175	164
	IB	NAD-2A	16.1	175	185
	IC	NAD-2A	16.1	175	183
	IIA	NAD-1	12.8	200	135
	IIB	NAD-1	12.8	200	195
	IIC	NAD-1	12.8	200	22 1
	IIIA	NAD-2	11.2	200	240
	IIIB	NAD-2	14.0	250	190
	IVA	NAD-3	14.0	225	150
	IVB	NAD-3	16.2	225	200
_	VIA	NAD-2	15.3	175	160 ·

^{*}Average of three readings

	EXAMPLE 14.	
5	Resin NAD-1 (110 parts) was formulated and blended to a #6 Hegman grind with 100 parts titanium dioxide, 0.015 parts cobalt drier (12% cobalt), 0.10 parts zirconium drier (12% zirconium) in three formulations A, B and C containing 5, 10 and 15 parts of tall oil alkyd (100% solids) respectively. Various weights of sodium hydroxide were then added as a 25 weight percent solution in methyl alcohol to form the solid paints identified in Table IV as Experiment IIA, IIB and IIC. Solid form the solid paints identified in Table 17 and 195 respectively exhibited	5
10	form the solid paints identified in Table IV as Experimental Paints IIA and IIB with gel strengths of 135 and 195 respectively exhibited satisfactory application characteristics. Solid paint IIC exhibited poor application characteristics. All three solid paints exhibited dimensional stability and gave a satisfactory dry coating on application to a test panel.	10
15	EXAMPLE 15. Resin NAD-2 (105 parts) was formulated and blended to a #6 Hegman grind with 100 parts titanium dioxide, 0.015 parts cobalt drier (12% cobalt), 0.10 parts zirconium drier (12% zirconium) and 10 parts tall oil alkyd (100%). Various weights of sodium hydroxide were added as a 25 weight percent solution in methyl alcohol to form the solid paints identified in Table IV as Experiments IIIA and IIIB. Solid paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics (too soft, paint IIIA with gel strength of 240 had inferior application characteristics).	15
20	paint IIIA with gel strength of 240 had interior applications of solid paint IIIB having a gel heavy drag) as opposed to the good characteristics of solid paint IIIB having a gel strength of 190. Although the paints exhibited dimensional stability the film appearance was poor due to unsatisfactory coalescence.	20
25	EXAMPLE 16. Resin NAD-3 (101 parts) was formulated as indicated for NAD-2 in Example 13 above using 10 parts tall oil in one case and replacing the tall oil with 15 parts of the polyester alkyd modifier in the second case. The corresponding solid paints prepared by the addition of a 25 weight percent solution of sodium hydroxide in methanol are identified in Table IV as solid paints IVA and IVB respectively. Solid paints IVA and IVB with gel strengths of 150 and 200 exhibited dimensional stability and satisfactory application and film characteristics.	25
30	EXAMPLE 17. Paint blocks of approximate size 4" × 6" formed from the above described solid paints were stored using a thin Saran (Trademark of the Dow Chemical solid paints were stored of six months. Application of these paints to a test	30
35	solid paints were stored using a thin Sarah (Hadelman of Company) envelope for a period of six months. Application of these paints to a test panel after the storage period showed no detectable deterioration of the application and film characteristics. Additionally solid paints prepared from the same resins but having acid values in the range of from 25 to 60 gave acceptable solid paint characteristics. The best application properties were obtained when the gel strength as measured by the Universal penetrometer was between 130 and 195 mm. although formulations having gel strength of from 100—130 and 195—200 gave effective solid paints with somewhat less desirable characteristics.	35
40	. TSCANDIE 10	
45	Resin NAD-4 (94 parts of 50 N/V suspension in mineral spirits) was formulated and blended to a #6 Hegman grind with 30 parts of alkyd modifier, 100 parts titanium dioxide, 15 parts calcium carbonate, 0.65 parts cobalt drier (12 percent cobalt), 0.65 parts manganese drier (8 percent metal), and 6.0 parts zirconium drier (12 percent zirconium). Sodium hydroxide (25% solution in methanol) was then added to form the solid paint identified in Table IV as experiment VI A. This solid paint had good application characteristics, exhibited dimensional stability and gave a dry film on a test panel.	45
50	EXAMPLE 19.	50
, JU	A 'non-aqueous dispersion' was prepared without using added stabilizer. A monomer system was chosen so that it would be partially swelled in the non-polar solvent, this being enough to maintain stability of the dispersion. In this case, 780 parts butyl acrylate, 100 parts methacrylic acid, 8 parts dodecyl mercaptan, 12 parts azobisisobutyronitrile, and 600 parts mineral spirits dodecyl mercaptan.	55
. 55	dodecyl mercaptan, 12 parts azobisisobutyrointrile, and doo held at 80°C. for 5 were charged to a reactor. The charge was brought to and held at 80°C. for 5 hours. Conversion was 97%, the acid value of the dispersion was 43.7. The theoretical acid value is 72, i.e. a certain amount of the acid is buried when this method of preparation is used.	

		,493,744		14
5	Two aliquots each of 180 parts parts (200% neutralization) and 37 hydroxide as a 25 percent solution dimensional stability; however, the a This product is not a true NAD dispersion. This does point out, how through a judicious choice of monom the ionizable sites are buried.	on in methanol. Both application characteristic and could best be described.	products exhibited s were poor. bed as a very coarse	5
. 10	PREPARATION OF NON-BONI		NON-AQUEOUS	10
15	no gelling sites) were prepared by the addition polymerization of various monomers in the presence of non-solvents, free radical initiators and stabilizers, one example of which is given in the table shown below. A small portion of the monomers is charged to the polymerization kettle with the non-solvent (mineral spirits etc.) and about 50 percent of the desired stabilizer and polymerization is initiated by heating to 75—80°C. After approximately 15—30 minutes, feeding of the remaining monomers, stabilizers, etc., is commenced and continued over 3—4			
	cooled to yield a milky dispersion have percent by weight non-volatile conter	ring a low viscosity (100- it.	-200 cps) and a 46	20
	Material .	Charge	Feed	
	ethyl acrylate	46	253	_
	methyl methacrylate	9	64	-
	odourless mineral spirits*	300	76	
	mineral spirits ^b	40	10	•
	stabilizer (38%)	37.5	37.5	
	azobis isobutyronitrile	1.5	4.0	
25	ALHOUSE 3 Wide variety of MAY	IPLE 21.		
	following illustrates the preparation stabilizer:	of a particularly desi	ig NAD resins the rable and useful	25
30	900 parts 12-hydroxystearic acid, 3. odourless mineral spirits were heated atmosphere. The reaction was monitore resulting product had an acid value of 3 reacted at 190°F, under nitrogen with parts triethylamine as a catalyst. This rea	d by collecting the by-process (calculated 33). The process of the calculated 33 is the process of the calculated 33 is the process of the calculated 33 is	oduct water. The oduct was further	30
35	parts triethylamine as a catalyst. This rea a non-volatile content of 91.0. This polymerized under free radical cond isobutyronitrile with 300 parts methyl m mineral spirits (800 parts). The produc mineral spirits. The final non-volatile co	second intermediate itions at 85°C using nethacrylate in the present	d value of 1.8 and (440 parts) was 2.0 parts azobis ence of odourless	35
40	Alkyd Resin J, having bonding sites paints in combination with the non-buspersions shown in Example 20, with the paints trimethylological pages 1295 parts trimethylological pages 1295 page	PLE 22. and particularly useful in ponding NAD Resin in as prepared by polyment	n preparing solid SON-AQUEOUS zing a mixture of	40
45	295 parts trimethylol propane, 690 parts of Azelaic dimer (AZELAIC 1110) and 423 250°C to an acid value of 42. The resulting.	achydrated castor on fath	y acid, 340 parts	45

EXAMPLE 23. Solid paints were prepared from Resin J using the following formulations:

Materials Pa	arts	
Experiment No.	1	2
	100	100
Non-Bonding NAD Resin	25	30
Ion-Bonded Resin J	5	
AC 100	•	1.2
Rheox I	1	
Titanium Dioxide	100	100
Min-U-Sil 10	. 10	10
Celite 499	10	10
Cobalt drier (12.0 percent metal)	0.65	0.70
	0.25	0.3
Manganese drier (9.0 percent metal)	2.5	3.0
Zirconium drier (12.0 percent metal)	2.0	1.0
Methanol	2.0	
Sodium hydroxide-methanol (24 percent sodium hydroxide)	6.0	7.0
	195	195
% neutralization	160	150
gel strength (mm.)		

The materials were added in the order shown, except that half of the quantity of The materials were added in the order shown, except that half of the quantity of NAD (50 parts) were held out until after the grind was achieved (i.e. all the pigments were added). Grind was 5½ hegman. After the remaining NAD was added and the batch cooled, the driers were added. The driers were allowed to mature for ½ hour before the sodium hydroxide-methanol was added with agitation under reduced pressure in a 'vacuum Cowles' to form a solid paint. This manner of addition diminishes the chances of entrapping air into the 'final' solid paint.

Both paints exhibited dimensional stability. When rubbed (by hand) on a substrate, paint was transferred to the substrate forming a film. Both films exhibited dry overnight.

The solid paints formed by combining Non-Bonding NAD resins with lesser quantities of ION-bonded resins of the IA type exemplified in Examples 1—12 and 5 5 10 10 quantities of ION-bonded resins of the IA type exemplified in Examples 1—12 and 15 quantities of 10IN-bonded resins of the IA type exemplified in Examples 1—12 and 22 are particularly for the trade sales (consumer) segment of the coating industry as well as for commercial coating applications such as for maintenance coatings and coil coating. The particular advantage of such combination and intercombination of ion-bonding and NAD resins (both bonding and bonding types) is that dimensional stability is retained with less bonding sites while application and film characteristics are greatly improved. 15 20 20

EXAMPLE 24.

The change in the proportion of Non-Bonding NAD resins shown in Example 23 from 75 to 200 parts NAD resir. per 25 parts of ion-bonded resin will yield equally satisfactory dimensionally stable solid paints.

The above Examples are illustrative of the best mode for the practice of this

invention and are not to be construed as limitations thereon.

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٠	WHAT WE CLAIM IS:— 1. A solid paint composition having a gel strength ranging from 100 to 200 millimeter penetration and a dimensional stability based on ion bonding comprising the admixture of:	
5	a. a polymer composition selected from: (1) a solution of an ion-bonding resin having a molecular weight ranging from 1,000 to 7,000 and sufficient reactive acid functional groups selected from carboxylic, sulfonic and phosphonic groups and provide an acid number from 20 to 80, said resin dissolved in a non-polar solvent to provide a 25 to 90 weight per cent	5
10	(2) a stabilized non-aqueous dispersion of a polymer having a molecular weight ranging from 25,000 to 1,000,000 and sufficient reactive acid functional groups selected from carboxylic, sulfonic, and phosphonic groups to provide an acid number from 25 to 60, said polymer dispersed as a 25 to 90 weight per cent	10
15	(3) a mixture of a non-bonding NAD resin comprising a stabilized dispersion of a polymer having a molecular weight ranging from 25,000 to 1,000,000 dispersed as a 25 to 90 weight per cent suspension in a non-polar non-solvent, said polymer having no reactive functional group sites with an ion-bonding regin solution or	15
20	defined in (1), the proportion by weight of non-bonding NAD resin to ion-bonding being from 2:1 to 8:1; and b. an ionic cross-linking agent selected from metal hydroxide, metal oxide ammonium hydroxide, metal alkoxide and an organic cation former dissolved or suspended an a polar solvent of high dielectric strength to provide a 10—50 weight	20
25	per cent solution or suspension; and c. a metallic drier in amounts from 0 to 5 weight per cent based on the total weight of polymer; wherein said composition contains from 100 to 600 mole per cent of ionic cross-linking agent per mole of acid functional group.	25
30	2. A composition according to claim I, wherein the cross-linking agent is a metal hydroxide selected from sodium, potassium, lithium, barium, calcium, manganese and magnesium hydroxides, the polar solvent is selected from the group consisting of a C_{1-8} aliphatic alcohol, formamide and water; and the functional group is a carboxylic acid group.	30
35	3. A composition according to claims 1 or 2, wherein the polar solvent is methanol and the cross-linking agent is sodium hydroxide. 4. A composition according to any one of claims 1 to 3, wherein the resin dissolved in a non-polar solvent is a polyester resin having a molecular weight of 1500—3500 and acid number 38—48 and is dissolved in mineral spirits and the ionizing cross-linking agent is codium that	35
40	ionizing cross-linking agent is sodium hydroxide present at 220—280 mole per cent excess basis the acid functional group. 5. A composition according to any one of claims 1 to 4, which comprises from 1 to 3 weight per cent of organic acid metal salt drier, said metal being selected from cobalt, zinc, magnesium, aluminum, manganese and zirconium.	40
45	6. A composition according to any one of claims 1 to 3, wherein the polymer composition comprises a non-aqueous dispersion of a resin having a molecular weight of 100,000 to 300,000 and acid number 30—40 suspended in mineral spirits and the sodium hydroxide is present at 120—280 mole per cent. 7. A composition according to claim 6, wherein the non-aqueous dispersion is formed from a virul acctate that the solice of the composition according to claim 6, wherein the non-aqueous dispersion is	45
50	formed from a vinyl acetate-ethyl acrylate methacrylic acid terpolymer resin having a molecular weight of 100,000 to 300,000 and an acid number of 25 to 80 is reacted with 210—250 mole per cent of sodium hydroxide as a 25 weight per cent solution in methanol. 8. A composition according to any one of the preceding claims, wherein the	50
55	dispersion with an ion-bonding resin solution, the proportion by weight of the former to the latter being from 2:1 to 4:1. 9. A process for prenaring a solid paint having dimensional stability based as	55
60	comprises: (a) dissolving or suspending a polymer having reactive acid functional groups according to claim 1 to form the respective polymer composition or mixtures thereof in such proportion to provide sufficient reactive acid functional groups.	60
65	necessary for the indicated dimensional stability when cross-linked by ionic cross-linking agents;	65

	(b) mixing pigments, fillers, colorants and 0 to 5 weight per cent of an organic acid metal salt drier into the resin solution;	
5	(c) adding thereto under vigorous stirring a 20 to 30 weight per cent solution of suspension of an ionic cross-linking agent selected from metal hydroxides, metal oxides, metal alkoxides, ammonium hydroxide and organic cation formers in a polar solvent at high dielectric strength containing 100 to 600 mole percent of the amount of ionic cross-linking agent required to neutralize said reactive acid groups	5
10	of the resin. 10. A process according to claim 9, wherein the resin of claim 1 (a) is a homopolymer selected from alkyd resin, polyester, unsaturated polyester, polymethacrylate and polyacrylate, or copolymers thereof with vinylchloride, an olefin or styrene, or mixtures thereof, wherein said reactive acid group is a	10
15	carboxylic acid group. 11. A process according to claim 10, wherein a polyester alkyd resin having a molecular weight of 1500 to 3500 and an acid number of 38 to 48 is reacted with 210—250 mole per cent of sodium hydroxide as a 25 weight per cent solution basis	15
20	mole of carboxylic acid group. 12. A process according to claim 9, wherein the resin is dispersed in the non- polar non-solvent at a content of 30 to 60 weight per cent in the presence of a stabilizer, said resin having a molecular weight of 25,000 to 1,000,000 and sufficiently pendant or terminal reactive acid functionality to yield an acid value in	20
25	the range of 25 to 60. 13. A process according to claims 10 to 12, wherein said reactive acid group is a carboxylic acid group present in amounts from 1 to 4 per 2,000 units of molecular weight. 14. A process according to claim 9, wherein the resin composition of claim 1	25
	(c) comprises a non-bonding non-aqueous dispersion with an ion-bonding reshifts solution in the ratio by weight of from 2:1 to 4:1. 15. A solid paint prepared according to the process of any one of claims 9 to	30
30	16. A paint stick which comprises the solid paint according to any one of claims 1 to 8, encased in a removable skin. 17. A solid paint composition according to claim 1, substantially as herein	3.
35	described with reference to the Examples. 18. A process for the preparation of a solid paint composition according to claim 9, substantially as herein described with reference to the Examples. 19. A composition according to claim 1, wherein the polymer composition is the ion-bonding resin according to a(1) or the stabilized non-aqueous dispersion	35
40	according to a(2). 20. A composition according to claim 1, wherein the polymer composition is the mixture of non-bonding NAD resin with the ion-bonding resin solution according to a(3).	40

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